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A peptide-based fluorescent probe images ERAAP activity in cells and in high throughput assays;

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ERAAP is an intracellular amino-peptidase that plays a central role in determining the repertoire of peptides displayed by cells by MHC class I molecules, and dysfunctions in ERAAP are linked to a variety of diseases. There is therefore great interest in developing probes that can image ERAAP in cells. In this report we present a fluorescent probe, termed Ep, that can image ERAAP activity in live cells. Ep is composed of a 10 amino acid ERAAP substrate that has a donor quencher pair conjugated to it, composed of BODIPY and dinitrotoluene. Ep undergoes a 20-fold increase in fluorescence after ERAAP cleavage, and was able to image ERAAP activity in cell culture via fluorescence microscopy. In addition, we used Ep to develop a high throughput screen for ERAAP inhibitors, and screened an electrophile library containing 1460 compounds. From this Ep based screen we identified aromatic alkyne-ketone as a lead fragment that can irreversibly inhibit ERAAP activity. We anticipate numerous applications of Ep given its unique ability to image ERAAP within cells.

ER aminopeptidase associated with antigen processing (ERAAP) is an intracellular amino-peptidase, located in the endoplasmic reticulum, which cleaves peptides presented by the major histocompatibility complex (MHC) class I molecules and determines which subset of peptides is displayed on cell surfaces and recognized by killer T cells.^{1–3} ERAAP thus plays a key role in shaping the immune response and its polymorphic variants have been implicated in a number of autoimmune diseases.^{4–6} Down-regulation of ERAAP results in alterations in the peptides that are displayed on cell surfaces. The altered peptide repertoire is recognized as foreign, because killer T cells are not tolerant towards the new peptides.^{7–9} Therapeutics that can down regulate ERAAP therefore have great potential as cancer vaccines and

increasing the immune response towards tumors.^{10,11} Downregulation of ERAAP is also implicated in the development of auto-immune disorders, such as ankylosing spondylitis and psoriasis.^{4,5}

There is therefore great interest in developing probes that can image ERAAP in cells and also in the development of new inhibitors for ERAAP. However, developing imaging probes for ERAAP has been challenging because it is an intracellular amino-peptidase that cleaves large peptides, and developing membrane permeable probes that can turn on fluorescence after ERAAP mediated cleavage remains a challenge. In addition, there are numerous other amino-peptidases within cells that will also potentially cleave fluorescent probes designed to specifically detect ERAAP. For example, the only fluorescent probe currently commercially available for measuring ERAAP activity is leucine-7amino-5-methylcoumarin (L-Amc), which is cleaved by all aminopeptidases, and also lacks the optimal emission wavelength for cellular imaging. Although recent progress has been made in developing ERAP1 (human ortholog of mouse ERAAP) imaging probes, using two photon emitting dyes,¹² there is still a great need for fluorescent probes that can detect ERAAP via conventional fluorescence microscopy.

In this report we present a fluorescent ERAAP probe, composed of a known ERAAP-substrate, the N-terminally extended KK-SIINFEKL decapeptide. The modified peptide has a donor quencher pair conjugated to it and is termed Ep. Ep emits at 510 nm after ERAAP cleavage, and can thus image ERAAP activity in cells. In addition, Ep was specific for ERAAP over other cellular amino-peptidases and has low activity in cells that lack ERAAP. Finally, Ep has high sensitivity for ERAAP and was used to develop a high-throughput-screen (HTS) for ERAAP inhibitors that allowed us to identify a novel electrophile fragment, alkyne-ketone that can inhibit ERAAP.

The chemical structure of Ep is shown in Fig. 1a, it is composed of the modified decapeptide K(DNP)-K-S-I-I-N-F-E-C(BODIPY)-L. Ep contains the fluorescent dye BODIPY and the fluorescent quencher dinitrophenyl (DNP), conjugated to the KK-SIINFECL peptide. Ep should initially have low fluorescence

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[†] Electronic supplementary information (ESI) available: Detailed procedures for Ep synthesis, kinetics, quantum yield, docking and Ep-based high throughput screening. See DOI: 10.1039/c7cc09598h



Fig. 1 Ep is a peptide-based fluorescent probe that can detect ERAAP. (a) Ep is based on the peptide sequence (KK-SIINFECL) and its fluorescence is quenched through a FRET pair between DNP and BODIPY. ERAAP cleaves the N-terminal amino acid of Ep and releases the quencher, causing a dramatic increase in fluorescence. (b) ERAAP activity imaged in wildtype fibroblasts and in ERAAP knockout fibroblasts. (c) A high throughput screening assay for identifying ERAAP inhibitors developed in a 96-well fluorescent plate reader format. A 1460-electrophile library was screened and we identified *para*-bromo-, *ortho*-fluoro-phenyl ketone alkyne as a lead electrophile fragment that can irreversibly inhibit ERAAP.

because the DNP group will quench the BODIPY fluorescence. However, in the presence of ERAAP the fluorescence of EP should increase, because ERAAP will cleave the N-terminal amino acid modified by the DNP group, due to its aminopeptidase activity. The diffusion of DNP away from the BODIPY should increase the quantum yield of Ep. The peptide KK-SIINFECL was selected as the ERAAP substrate for Ep because of its similarity to the well characterized ERAAP substrate KK-SIINFEKL, which is trimmed by ERAAP to SIINFEKL, the immunodominant H-2Kbbinding peptide.^{3,13} SIINFECL is identical to SIINFEKL except that its lysine residue is replaced with a cysteine, which was needed for conjugation to BODIPY. Importantly, Serwold et al. demonstrated that the internal lysine in SIINFEKL containing peptides can be modified to other residues without affecting its ability to bind the Kb MHC I molecule.14,15 Based on this design, Ep was synthesized in one step using the commercially synthesized peptide K(DNP)K-SIINFECL, which contained the fluorescent quencher, DNP (dinitrophenyl).16-18 The BODIPY (4,4-difluoro-3a,4a-diaza-s-indacene), a widely used fluorophore for fluorescence imaging in life sciences, was conjugated to this peptide via a reaction between the cysteine side-chain and iodoacetylamide BODIPY (see the ESI† for the reaction scheme



and procedures). The final Ep was purified by reverse-phase HPLC and its synthesis was confirmed by ESI-HRMS (see Fig. S1, ESI[†]).

Ep is designed to be initially non-fluorescent because of DNP mediated quenching, but should increase its fluorescence after ERAAP cleavage. We therefore investigated the fluorescence of Ep in its intact state, and its fluorescence after cleavage by Ep. The quantum yield of Ep is extremely low in aqueous solution, about 1%, due to efficient fluorescence quenching. However, upon addition of ERAAP the fluorescence quantum yield of Ep dramatically increases, by a factor of 22 due to ERAAP mediated cleavage, resulting in bright green fluorescence (see Fig. S2 in the ESI†). Fig. 2 demonstrates that the Ep can sensitively detect ERAAP and was able to detect ERAAP at concentrations as low as $0.1 \,\mu g \, \text{mL}^{-1}$, and quasi-linearly responded to ERAAP concentrations within the sub $\mu g \, \text{mL}^{-1}$ range.

We performed HPLC experiments to determine if Ep was sequentially trimmed by ERAAP at the N terminus to release the DNP residue. Ep was incubated with ERAAP and HPLC was performed in the reaction mixture, after 10 min, and the absorption at 350 nm (DNP) and 490 nm (BODIPY) was measured. Fig. 3 demonstrates that Ep by itself gives only one peak at 350 nm and 490 nm, which overlap. However, after incubation with ERAAP, the original peak decreased and several new peaks emerged corresponding to the products of the enzymatic reaction. We studied the kinetics of the enzymatic trimming reaction between Ep and ERAAP using fluorogenic assay, by plotting the initial reaction velocities against Ep concentrations. Ep underwent



Fig. 2 Ep can detect ERAAP activity. The fluorescence intensity of Ep (at 510 nm) changes in response to different concentrations of ERAAP after incubation in $1 \times$ PBS for 10 min. (Inset: Response curve in the range below 1 µg mL⁻¹ of ERAAP).

Fig. 3 HPLC analysis of Ep before and after trimming by ERAAP.

typical Michaelis–Menten hydrolysis kinetics (see Fig. S3 in the ESI†), and fitting of the kinetic data to a Lineweaver–Burk plot gave the kinetics parameters $K_{\rm m} = 6.7 \,\mu$ M, $V_{\rm max} = 1.14 \,$ nM s⁻¹, and $k_{\rm cat} = V_{\rm max}/[\text{ERAAP}] = 0.13 \,$ s⁻¹, corresponding to a catalytic efficiency ($k_{\rm cat}/K_{\rm m}$) = 1.94 × 10⁴ M⁻¹ s⁻¹. The $K_{\rm m}$ and $k_{\rm cat}$ of Ep for ERAAP are comparable to other known substrates of ERAAP, and suggests that it will be able to compete with endogenous cellular substrates for ERAAP in cells.¹⁹

To further understand the binding mechanism of Ep with ERAAP, we performed molecular docking of Ep with ERAAP, using the solved crystal structure of ERAP1 (human ortholog of mouse ERAAP, PDB 2DY0). ERAP1 has a large and long binding pocket that contains two critical binding sites at its terminal two ends. The Zn-associated catalytic site is at one end of the binding pocket and binds the N-terminus of the target peptide and performs the trimming. The other peptide binding site is the regulatory site (\sim 30 Å away from the catalytic center) which binds the C-terminus of the peptide and could function as a molecular ruler to control the trimming selectivity. The middle part of the binding pocket has a large volume and high conformational flexibility.¹⁹⁻²²

We decided to dock only the first 4 and last 3 amino acid residues of Ep (tetrapeptide and tripeptide) to the active site and regulatory site, respectively, due to the challenges associated with docking a large peptide to a flexible pocket in a protein. Fig. 4a shows the optimized conformation of the tetrapeptide, K(DNP)KSI, in the catalytic site. We found that the N-terminal amine group and the first amide bond were located close to the Zn center and the carbonyl group was stabilized via hydrogen bonding with Tyr438 and His353. The bulky DNP side chain of N-terminal Lys(DNP) fell into the hydrophobic sub-pocket S1 through hydrophobic interactions, the rest of the peptide chain extended backward from the catalytic site. We compared this docking result with the crystal structure of the ERAP1-Bestatin complex (PDB 2DY0). Bestatin is a widely used Zn metallo-aminopeptidase inhibitor which tightly binds into the catalytic site and we found that the tetrapeptide K(DNP)KSI and Bestatin shared very similar conformations in the catalytic site (Fig. 4a and b). The results above

demonstrated that the N-terminus of Ep has great affinity to the ERAP1 catalytic site, which triggers the trimming process. The docking result of the C-terminal tripeptide EC(BODIPY)L with the ERAP1 regulatory site is shown in Fig. 4c. The carboxylate end of the tripeptide was strongly anchored in the ERAP1 regulatory site by direct and water mediated hydrogen bonds with Tyr684, Lys685, Arg807 and Arg841, and the BODIPY residue on the side chain of the cysteine fell into the non-polar domain surrounded by Leu733 and Leu734. This result indicates that the C-terminus of Ep can form strong binding with the ERAP1 regulatory site. It is also in agreement with the reported binding preference of the ERAP1 regulatory site towards hydrophobic side chains.²⁰

There is great interest in being able to image ERAAP activity in cells and we therefore investigated if Ep could allow this. For these experiments we incubated Ep with a fibroblast cell line generated from B6 wildtype (WT) mouse and from an ERAAP knockout (ERAAP-KO) mouse. We treated WT and ERAAP-KO fibroblast cells with 10 or 20 μ M of Ep for 2 h or 3 h, and then imaged *via* fluorescence microscopy. Fig. 5 demonstrates that Ep can image ERAAP activity in live cells. For example, strong fluorescence was detected in WT cells, in contrast, ERAAP-KO cells did not show any observable fluorescence signal, in all Ep concentrations and incubation times investigated. These results demonstrate that Ep can detect ERAAP in cells in a sensitive and specific manner.

The downregulation of ERAP1 (human ortholog of ERAAP) triggers potent anticancer immune responses and can rescue mice from experimental models of cancer. Therefore, there is great interest in developing ERAAP inhibitors for cancer immunotherapy.^{10,11} However, developing ERAAP inhibitors has been challenging because of the lack of assays available for detecting ERAAP with sensitivity in a high throughput manner. Current ERAAP assays have low selectivity and sensitivity, and are therefore challenging to use for the development of HTS assays for identifying ERAAP (ERAP1) inhibitors. We therefore investigated if Ep had the sensitivity needed to develop an HTS assay for identifying ERAAP inhibitors. We determined the Z factor of Ep, with regard to developing ERAAP inhibitors, by measuring its fluorescence in the presence and in the absence of ERAAP. In a 96-well plate format, this assay gave a Z'-factor of 0.79, which demonstrates that it is suitable for performing HTS screens



Fig. 4 (a) Optimized confirmations of the tetrapeptide (K(DNP)KSI) in the ERAP1 catalytic site. (b) Crystal structure of Bestatin in the ERAP1 catalytic site (PDB: 2DY0). (c) Optimized confirmations of the tripeptide EC(BODIPY)L in the ERAP1 regulatory site.



Fig. 5 Fluorescence imaging of wildtype (WT) and ERAAP knockout (ERAAP-KO) fibroblast cells with Ep. WT cells generate green fluorescence after Ep incubation, whereas ERAAP-KO fibroblasts generate minimal fluorescence.



Fig. 6 Ep based high throughput screen identifies electrophiles that can inhibit ERAAP activity. (a) Scatter plot describing the results of the 1460-compound screen for inhibition of ERAAP activity. (b) Validation of the top 8 hits for ERAAP inhibition. Compounds **A**, **B** and **C** showed significantly higher inhibition efficiency (\geq 62%) than the other 5 hit compounds. (c) Dose-dependent inhibition curves of compounds **A**, **B** and **C**. Compound **A** had an IC₅₀ of 23 µM, which was the lowest among all three compounds. (d) Structure of compound **A**, a promising pharmacophore for developing irreversible ERAAP inhibitors.

(see Fig. S5 in the ESI[†]). Ep was used to screen a 1460- compound electrophile library (Enamine Ltd). We chose an electrophile library for screening because these fragments can serve as leads for the development of irreversible inhibitors.

The results of our HTS are shown in Fig. 6 as an activity scatter-plot (Fig. 6a). We identified 8 compounds that could reproducibly inhibit ERAAP activity. Three compounds (A-C) were chosen for further study because of their low IC₅₀ values in comparison to the other members of the library (Fig. 6b). Of these, compound A, (1-(4-bromo-2-fluorophenyl)prop)-2-yn-1one, exhibited the lowest IC₅₀ (23 μ M), whereas compounds B and C had IC₅₀ values of 92 µM and 175 µM, respectively (Fig. 6c). Although this value is still high compared with several other reported ERAP1 inhibitors, which have sub-micromolar range $IC_{50}s$,^{23,24} we anticipate that the efficiency of A-C can considerably be improved after further optimization. Steadystate time dependent inhibition studies with A demonstrated that it could generate almost 100% of inhibition of ERAAP after a 4.5 h pre-incubation (see Fig. S6 in the ESI⁺), suggesting that it is an irreversible inhibitor. Compound A is a halogen substituted phenyl ketone alkyne (Fig. 6d), and its small size and low electrophilicity suggest that it is a suitable pharmacophore for developing irreversible ERAAP inhibitors.

In conclusion, in this report we present a new peptide based fluorescent probe for ERAAP, termed Ep, which can image ERAAP activity in cells and in a plate reader format. We anticipate numerous applications for Ep given the great interest in understanding the biology of ERAAP and in developing new ERAAP inhibitors.

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Conflicts of interest

There are no conflicts to declare.

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